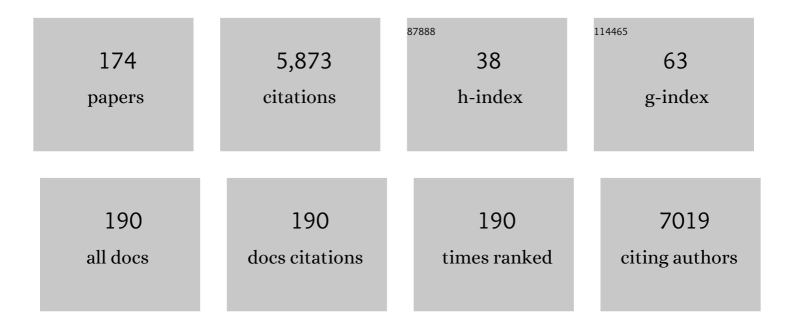
Mingyue Zheng

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Facing small and biased data dilemma in drug discovery with enhanced federated learning approaches. Science China Life Sciences, 2022, 65, 529-539.	4.9	17
2	New <scp>Cembraneâ€Type</scp> Diterpenoids from the South China Sea Soft Coral <i>Sinularia nanolobata</i> . Chinese Journal of Chemistry, 2022, 40, 28-38.	4.9	10
3	Multi-instance learning of graph neural networks for aqueous p <i>K</i> a prediction. Bioinformatics, 2022, 38, 792-798.	4.1	19
4	Drug target inference by mining transcriptional data using a novel graph convolutional network framework. Protein and Cell, 2022, 13, 281-301.	11.0	18
5	Discovery of Pyrazolo[3,4- <i>d</i>]pyridazinone Derivatives as Selective DDR1 Inhibitors via Deep Learning Based Design, Synthesis, and Biological Evaluation. Journal of Medicinal Chemistry, 2022, 65, 103-119.	6.4	31
6	Graph neural network approaches for drug-target interactions. Current Opinion in Structural Biology, 2022, 73, 102327.	5.7	51
7	Optimization of metabolomic data processing using NOREVA. Nature Protocols, 2022, 17, 129-151.	12.0	114
8	Discovery of ARF1-targeting inhibitor demethylzeylasteral as a potential agent against breast cancer. Acta Pharmaceutica Sinica B, 2022, 12, 2619-2622.	12.0	5
9	Discovery of the First-in-Class Agonist-Based SOS1 PROTACs Effective in Human Cancer Cells Harboring Various KRAS Mutations. Journal of Medicinal Chemistry, 2022, 65, 3923-3942.	6.4	36
10	An inductive graph neural network model for compound–protein interaction prediction based on a homogeneous graph. Briefings in Bioinformatics, 2022, 23, .	6.5	9
11	Bloodâ \in "brain barrier penetration prediction enhanced by uncertainty estimation. Journal of Cheminformatics, 2022, 14, .	6.1	11
12	Design, synthesis and biological evaluation of pyrazolo[3,4-d]pyridazinone derivatives as covalent FGFR inhibitors. Acta Pharmaceutica Sinica B, 2021, 11, 781-794.	12.0	16
13	DrugSpaceX: a large screenable and synthetically tractable database extending drug space. Nucleic Acids Research, 2021, 49, D1170-D1178.	14.5	23
14	Ligand-Promoted Alkynylation of Aryl Ketones: A Practical Tool for Structural Diversity in Drugs and Natural Products. ACS Catalysis, 2021, 11, 1758-1764.	11.2	28
15	Diversified strategy for the synthesis of DNA-encoded oxindole libraries. Chemical Science, 2021, 12, 2841-2847.	7.4	32
16	Drug repurposing against breast cancer by integrating drug-exposure expression profiles and drug–drug links based on graph neural network. Bioinformatics, 2021, 37, 2930-2937.	4.1	25
17	Cytoplasmic DNA sensing by KU complex in aged CD4+ TÂcell potentiates TÂcell activation and aging-related autoimmune inflammation. Immunity, 2021, 54, 632-647.e9.	14.3	37
18	Graph neural networks for automated de novo drug design. Drug Discovery Today, 2021, 26, 1382-1393.	6.4	71

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19	Discovery of a potent, selective, and covalent ZAP-70 kinase inhibitor. European Journal of Medicinal Chemistry, 2021, 219, 113393.	5.5	5
20	Targeting sorting nexin 10 improves mouse colitis via inhibiting PIKfyve-mediated TBK1/c-Rel signaling activation. Pharmacological Research, 2021, 169, 105679.	7.1	3
21	Generative Models for De Novo Drug Design. Journal of Medicinal Chemistry, 2021, 64, 14011-14027.	6.4	72
22	Crowdsourced identification of multi-target kinase inhibitors for RET- and TAU- based disease: The Multi-Targeting Drug DREAM Challenge. PLoS Computational Biology, 2021, 17, e1009302.	3.2	7
23	Structure-Based Virtual Screening and Identification of Potential Inhibitors of SARS-CoV-2 S-RBD and ACE2 Interaction. Frontiers in Chemistry, 2021, 9, 740702.	3.6	15
24	A hybrid framework for improving uncertainty quantification in deep learning-based QSAR regression modeling. Journal of Cheminformatics, 2021, 13, 69.	6.1	18
25	DFT Mechanism of Cu Catalyzed Coupling Reaction to Alkyl Aryl Ethers. Acta Chimica Sinica, 2021, 79, 948.	1.4	1
26	Regulating Glucose Metabolism with Prodrug Nanoparticles for Promoting Photoimmunotherapy of Pancreatic Cancer. Advanced Science, 2021, 8, 2002746.	11.2	96
27	SNX10â€mediated LPS sensing causes intestinal barrier dysfunction via a caspaseâ€5â€dependent signaling cascade. EMBO Journal, 2021, 40, e108080.	7.8	22
28	Active Learning for Drug Design: A Case Study on the Plasma Exposure of Orally Administered Drugs. Journal of Medicinal Chemistry, 2021, 64, 16838-16853.	6.4	12
29	AutoGGN: A gene graph network AutoML tool for multi-omics research. Artificial Intelligence in the Life Sciences, 2021, 1, 100019.	2.2	4
30	Current status of active learning for drug discovery. Artificial Intelligence in the Life Sciences, 2021, 1, 100023.	2.2	3
31	Discovery and characterization of natural products as novel indoleamine 2,3-dioxygenase 1 inhibitors through high-throughput screening. Acta Pharmacologica Sinica, 2020, 41, 423-431.	6.1	6
32	SNX10 (sorting nexin 10) inhibits colorectal cancer initiation and progression by controlling autophagic degradation of SRC. Autophagy, 2020, 16, 735-749.	9.1	43
33	Pushing the Boundaries of Molecular Representation for Drug Discovery with the Graph Attention Mechanism. Journal of Medicinal Chemistry, 2020, 63, 8749-8760.	6.4	402
34	Deep Learning Enhancing Kinome-Wide Polypharmacology Profiling: Model Construction and Experiment Validation. Journal of Medicinal Chemistry, 2020, 63, 8723-8737.	6.4	58
35	Discovery of Highly Potent, Selective, and Orally Efficacious p300/CBP Histone Acetyltransferases Inhibitors. Journal of Medicinal Chemistry, 2020, 63, 1337-1360.	6.4	85
36	Automated design and optimization of multitarget schizophrenia drug candidates by deep learning. European Journal of Medicinal Chemistry, 2020, 204, 112572.	5.5	25

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37	Characterization of covalent binding of tyrosine kinase inhibitors to plasma proteins. Drug Metabolism and Pharmacokinetics, 2020, 35, 456-465.	2.2	12
38	Sorting Nexin 10 Mediates Metabolic Reprogramming of Macrophages in Atherosclerosis Through the Lyn-Dependent TFEB Signaling Pathway. Circulation Research, 2020, 127, 534-549.	4.5	32
39	Revisiting Aldehyde Oxidase Mediated Metabolism in Drug-like Molecules: An Improved Computational Model. Journal of Medicinal Chemistry, 2020, 63, 6523-6537.	6.4	10
40	Solution-Phase DNA-Compatible Pictet-Spengler Reaction Aided by Machine Learning Building Block Filtering. IScience, 2020, 23, 101142.	4.1	13
41	Optimizing chemical reaction conditions using deep learning: a case study for the Suzuki–Miyaura cross-coupling reaction. Organic Chemistry Frontiers, 2020, 7, 2269-2277.	4.5	21
42	Discovery of novel glyceraldehyde-3-phosphate dehydrogenase inhibitor via docking-based virtual screening. Bioorganic Chemistry, 2020, 96, 103620.	4.1	17
43	Investigation of the remote acyl group participation in glycosylation from conformational perspectives by using trichloroacetimidate as the acetyl surrogate. Organic Chemistry Frontiers, 2020, 7, 1606-1615.	4.5	19
44	TransformerCPI: improving compound–protein interaction prediction by sequence-based deep learning with self-attention mechanism and label reversal experiments. Bioinformatics, 2020, 36, 4406-4414.	4.1	190
45	The application of artificial intelligence to drug sensitivity prediction. Chinese Science Bulletin, 2020, 65, 3551-3561.	0.7	Ο
46	Discovery of triazoloquinoxaline as novel STING agonists via structure-based virtual screening. Bioorganic Chemistry, 2020, 100, 103958.	4.1	6
47	Bioactivity Prediction Based on Matched Molecular Pair and Matched Molecular Series Methods. Current Pharmaceutical Design, 2020, 26, 4195-4205.	1.9	1
48	Improving the Virtual Screening Ability of Target-Specific Scoring Functions Using Deep Learning Methods. Frontiers in Pharmacology, 2019, 10, 924.	3.5	34
49	Discovery and Development of a Series of Pyrazolo[3,4- <i>d</i>]pyridazinone Compounds as the Novel Covalent Fibroblast Growth Factor Receptor Inhibitors by the Rational Drug Design. Journal of Medicinal Chemistry, 2019, 62, 7473-7488.	6.4	28
50	KinomeX: a web application for predicting kinome-wide polypharmacology effect of small molecules. Bioinformatics, 2019, 35, 5354-5356.	4.1	24
51	Deep Neural Network Classifier for Virtual Screening Inhibitors of (S)-Adenosyl-L-Methionine (SAM)-Dependent Methyltransferase Family. Frontiers in Chemistry, 2019, 7, 324.	3.6	10
52	Metabolomics Analysis of L-Arginine Induced Gastrointestinal Motility Disorder in Rats Using UPLC-MS After Magnolol Treatment. Frontiers in Pharmacology, 2019, 10, 183.	3.5	18
53	Hydrolytic Metabolism of Cyanopyrrolidine DPP-4 Inhibitors Mediated by Dipeptidyl Peptidases. Drug Metabolism and Disposition, 2019, 47, 238-248.	3.3	6
54	Rational design of 5-((1H-imidazol-1-yl)methyl)quinolin-8-ol derivatives as novel bromodomain-containing protein 4 inhibitors. European Journal of Medicinal Chemistry, 2019, 163, 281-294.	5.5	13

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55	Preclinical characterization of anlotinib, a highly potent and selective vascular endothelial growth factor receptorâ€2 inhibitor. Cancer Science, 2018, 109, 1207-1219.	3.9	233
56	Computational chemical biology and drug design: Facilitating protein structure, function, and modulation studies. Medicinal Research Reviews, 2018, 38, 914-950.	10.5	38
57	Machine Learning-Based Modeling of Drug Toxicity. Methods in Molecular Biology, 2018, 1754, 247-264.	0.9	14
58	Development and evaluation of a novel series of Nitroxoline-derived BET inhibitors with antitumor activity in renal cell carcinoma. Oncogenesis, 2018, 7, 83.	4.9	10
59	Discovery of Novel Inhibitors of Indoleamine 2,3-Dioxygenase 1 Through Structure-Based Virtual Screening. Frontiers in Pharmacology, 2018, 9, 277.	3.5	21
60	Artificial intelligence in drug design. Science China Life Sciences, 2018, 61, 1191-1204.	4.9	145
61	Predicting Hepatotoxicity of Drug Metabolites Via an Ensemble Approach Based on Support Vector Machine. Combinatorial Chemistry and High Throughput Screening, 2018, 20, 839-849.	1.1	4
62	Analysis and prediction of drug–drug interaction by minimum redundancy maximum relevance and incremental feature selection. Journal of Biomolecular Structure and Dynamics, 2017, 35, 312-329.	3.5	81
63	Discovery of novel BRD4 inhibitors by high-throughput screening, crystallography, and cell-based assays. Bioorganic and Medicinal Chemistry Letters, 2017, 27, 2003-2009.	2.2	9
64	Discovery of Novel Disruptor of Silencing Telomeric 1-Like (DOT1L) Inhibitors using a Target-Specific Scoring Function for the (<i>S</i>)-Adenosyl- <scp> </scp> -methionine (SAM)-Dependent Methyltransferase Family. Journal of Medicinal Chemistry, 2017, 60, 2026-2036.	6.4	22
65	Aldehyde Oxidase Mediated Metabolism in Drug-like Molecules: A Combined Computational and Experimental Study. Journal of Medicinal Chemistry, 2017, 60, 2973-2982.	6.4	34
66	Machine-Learning-Assisted Approach for Discovering Novel Inhibitors Targeting Bromodomain-Containing Protein 4. Journal of Chemical Information and Modeling, 2017, 57, 1677-1690.	5.4	40
67	Design, Synthesis, and Pharmacological Evaluation of Novel Multisubstituted Pyridin-3-amine Derivatives as Multitargeted Protein Kinase Inhibitors for the Treatment of Non-Small Cell Lung Cancer. Journal of Medicinal Chemistry, 2017, 60, 6018-6035.	6.4	19
68	Discovery and optimization of selective inhibitors of protein arginine methyltransferase 5 by docking-based virtual screening. Organic and Biomolecular Chemistry, 2017, 15, 3648-3661.	2.8	28
69	Discovery of novel trimethoxy-ring BRD4 bromodomain inhibitors: AlphaScreen assay, crystallography and cell-based assay. MedChemComm, 2017, 8, 1322-1331.	3.4	13
70	Discovery of novel BET inhibitors by drug repurposing of nitroxoline and its analogues. Organic and Biomolecular Chemistry, 2017, 15, 9352-9361.	2.8	34
71	Conformation and dynamics of the C-terminal region in human phosphoglycerate mutase 1. Acta Pharmacologica Sinica, 2017, 38, 1673-1682.	6.1	9
72	Aspirin Inhibits Cancer Metastasis and Angiogenesis via Targeting Heparanase. Clinical Cancer Research, 2017, 23, 6267-6278.	7.0	94

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73	RD-Metabolizer: an integrated and reaction types extensive approach to predict metabolic sites and metabolites of drug-like molecules. Chemistry Central Journal, 2017, 11, 65.	2.6	5
74	Structure-activity relationships of the antimalarial agent artemisinin and the research progress on the artemisinin analogues with novel pharmacological actions. Chinese Science Bulletin, 2017, 62, 1948-1963.	0.7	2
75	Pharmacologic characterization of SHR8443, a novel dual inhibitor of phosphatidylinositol 3-kinase and mammalian target of rapamycin. Oncotarget, 2017, 8, 107977-107990.	1.8	6
76	In Silico Prediction of Chemical Toxicity Profile Using Local Lazy Learning. Combinatorial Chemistry and High Throughput Screening, 2017, 20, 346-353.	1.1	10
77	Identification of Drug-Drug Interactions Using Chemical Interactions. Current Bioinformatics, 2017, 12, .	1.5	92
78	Integrated Analysis of Multiscale Large-Scale Biological Data for Investigating Human Disease 2016. BioMed Research International, 2016, 2016, 1-2.	1.9	2
79	Identification of novel candidate drivers connecting different dysfunctional levels for lung adenocarcinoma using protein-protein interactions and a shortest path approach. Scientific Reports, 2016, 6, 29849.	3.3	28
80	Discovery of a new series of imidazo[1,2-a]pyridine compounds as selective c-Met inhibitors. Acta Pharmacologica Sinica, 2016, 37, 698-707.	6.1	22
81	Estimation of elimination half-lives of organic chemicals in humans using gradient boosting machine. Biochimica Et Biophysica Acta - General Subjects, 2016, 1860, 2664-2671.	2.4	27
82	Discovery of Novel Inhibitors Targeting the Menin-Mixed Lineage Leukemia Interface Using Pharmacophore- and Docking-Based Virtual Screening. Journal of Chemical Information and Modeling, 2016, 56, 1847-1855.	5.4	22
83	ldentification of compound–protein interactions through the analysis of gene ontology, KEGG enrichment for proteins and molecular fragments of compounds. Molecular Genetics and Genomics, 2016, 291, 2065-2079.	2.1	62
84	Applying high-performance computing in drug discovery and molecular simulation. National Science Review, 2016, 3, 49-63.	9.5	30
85	Structural optimization and biological evaluation of 1,5-disubstituted pyrazole-3-carboxamines as potent inhibitors of human 5-lipoxygenase. Acta Pharmaceutica Sinica B, 2016, 6, 32-45.	12.0	11
86	Wedelolactone metabolism in rats through regioselective glucuronidation catalyzed by uridine diphosphate-glucuronosyltransferases 1As (UGT1As). Phytomedicine, 2016, 23, 340-349.	5.3	11
87	Identification of new candidate drugs for lung cancer using chemical–chemical interactions, chemical–protein interactions and a K-means clustering algorithm. Journal of Biomolecular Structure and Dynamics, 2016, 34, 906-917.	3.5	30
88	Identification of Novel Disruptor of Telomeric Silencing 1-like (DOT1L) Inhibitors through Structure-Based Virtual Screening and Biological Assays. Journal of Chemical Information and Modeling, 2016, 56, 527-534.	5.4	27
89	Combinatorial Pharmacophore Modeling of Multidrug and Toxin Extrusion Transporter 1 Inhibitors: a Theoretical Perspective for Understanding Multiple Inhibitory Mechanisms. Scientific Reports, 2015, 5, 13684.	3.3	15
90	<i>In silico</i> ADME/T modelling for rational drug design. Quarterly Reviews of Biophysics, 2015, 48, 488-515.	5.7	250

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91	Integrated Analysis of Multiscale Large-Scale Biological Data for Investigating Human Disease. BioMed Research International, 2015, 2015, 1-2.	1.9	0
92	Combinatorial Pharmacophore-Based 3D-QSAR Analysis and Virtual Screening of FGFR1 Inhibitors. International Journal of Molecular Sciences, 2015, 16, 13407-13426.	4.1	26
93	Prediction of Drug Indications Based on Chemical Interactions and Chemical Similarities. BioMed Research International, 2015, 2015, 1-14.	1.9	3
94	A Genetic Algorithm Based Support Vector Machine Model for Blood-Brain Barrier Penetration Prediction. BioMed Research International, 2015, 2015, 1-13.	1.9	23
95	TarPred: a web application for predicting therapeutic and side effect targets of chemical compounds. Bioinformatics, 2015, 31, 2049-2051.	4.1	52
96	Design, synthesis and biological evaluation of isoquinoline-based derivatives as novel histone deacetylase inhibitors. Bioorganic and Medicinal Chemistry, 2015, 23, 5881-5890.	3.0	17
97	Discovery and Optimization of Novel, Selective Histone Methyltransferase SET7 Inhibitors by Pharmacophore- and Docking-Based Virtual Screening. Journal of Medicinal Chemistry, 2015, 58, 8166-8181.	6.4	59
98	The Use of Chemical-Chemical Interaction and Chemical Structure to Identify New Candidate Chemicals Related to Lung Cancer. PLoS ONE, 2015, 10, e0128696.	2.5	9
99	Prediction of Cancer Drugs by Chemical-Chemical Interactions. PLoS ONE, 2014, 9, e87791.	2.5	14
100	Novel Bayesian classification models for predicting compounds blocking hERG potassium channels. Acta Pharmacologica Sinica, 2014, 35, 1093-1102.	6.1	53
101	Cholesterolâ€Î² ₁ AR interaction versus cholesterolâ€Î² ₂ AR interaction. Proteins: Structure, Function and Bioinformatics, 2014, 82, 760-770.	2.6	29
102	Metadynamics Simulation Study on the Conformational Transformation of Hhal Methyltransferase: An Induced-Fit Base-Flipping Hypothesis. BioMed Research International, 2014, 2014, 1-13.	1.9	14
103	In Silicotarget fishing: addressing a "Big Data―problem by ligand-based similarity rankings with data fusion. Journal of Cheminformatics, 2014, 6, 33.	6.1	48
104	Characterizing the binding of annexin V to a lipid bilayer using molecular dynamics simulations. Proteins: Structure, Function and Bioinformatics, 2014, 82, 312-322.	2.6	21
105	Rapid Generation of Privileged Substructure-Based Compound Libraries with Structural Diversity and Drug-Likeness. ACS Combinatorial Science, 2014, 16, 184-191.	3.8	30
106	Virtual screening and biological evaluation of novel small molecular inhibitors against protein arginine methyltransferase 1 (PRMT1). Organic and Biomolecular Chemistry, 2014, 12, 9665-9673.	2.8	27
107	Astemizole Arrests the Proliferation of Cancer Cells by Disrupting the EZH2-EED Interaction of Polycomb Repressive Complex 2. Journal of Medicinal Chemistry, 2014, 57, 9512-9521.	6.4	96
108	Repair of methyl lesions in RNA by oxidative demethylation. MedChemComm, 2014, 5, 1797-1803.	3.4	7

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109	<i>In silico</i> site of metabolism prediction for human UCT-catalyzed reactions. Bioinformatics, 2014, 30, 398-405.	4.1	29
110	Identifying Novel Selective Non-Nucleoside DNA Methyltransferase 1 Inhibitors through Docking-Based Virtual Screening. Journal of Medicinal Chemistry, 2014, 57, 9028-9041.	6.4	96
111	Catalytic Mechanism of Histone Acetyltransferase p300: From the Proton Transfer to Acetylation Reaction. Journal of Physical Chemistry B, 2014, 118, 2009-2019.	2.6	28
112	Estimation of acute oral toxicity in rat using local lazy learning. Journal of Cheminformatics, 2014, 6, 26.	6.1	30
113	Mechanism of the All- \hat{l} t o All- \hat{l}^2 Conformational Transition of RfaH-CTD: Molecular Dynamics Simulation and Markov State Model. Journal of Chemical Theory and Computation, 2014, 10, 2255-2264.	5.3	37
114	Interaction Between DNA/histone Methyltransferases and their Inhibitors. Current Medicinal Chemistry, 2014, 22, 360-372.	2.4	10
115	A quantum mechanics/molecular mechanics study on the hydrolysis mechanism of New Delhi metallo-β-lactamase-1. Journal of Computer-Aided Molecular Design, 2013, 27, 247-256.	2.9	36
116	Computational methods for drug design and discovery: focus on China. Trends in Pharmacological Sciences, 2013, 34, 549-559.	8.7	70
117	Identification of Novel Small Molecules as Inhibitors of Hepatitis C Virus by Structure-Based Virtual Screening. International Journal of Molecular Sciences, 2013, 14, 22845-22856.	4.1	12
118	Structure-Based Design and Synthesis of C-1- and C-4-Modified Analogs of Zanamivir as Neuraminidase Inhibitors. Journal of Medicinal Chemistry, 2013, 56, 671-684.	6.4	50
119	Water PMF for predicting the properties of water molecules in protein binding site. Journal of Computational Chemistry, 2013, 34, 583-592.	3.3	25
120	Binding sensitivity of adefovir to the polymerase from different genotypes of HBV: molecular modeling, docking and dynamics simulation studies. Acta Pharmacologica Sinica, 2013, 34, 319-328.	6.1	9
121	Rapid and selective access to three distinct sets of indole-based heterocycles from a single set of Ugi-adducts under microwave heating. Chemical Communications, 2013, 49, 2894.	4.1	48
122	Anthraquinone Derivatives as Potent Inhibitors of c-Met Kinase and the Extracellular Signaling Pathway. ACS Medicinal Chemistry Letters, 2013, 4, 408-413.	2.8	12
123	Combinatorial Pharmacophore Modeling of Organic Cation Transporter 2 (OCT2) Inhibitors: Insights into Multiple Inhibitory Mechanisms. Molecular Pharmaceutics, 2013, 10, 4611-4619.	4.6	21
124	Predicting Drugs Side Effects Based on Chemical-Chemical Interactions and Protein-Chemical Interactions. BioMed Research International, 2013, 2013, 1-8.	1.9	23
125	Prediction of Effective Drug Combinations by Chemical Interaction, Protein Interaction and Target Enrichment of KEGC Pathways. BioMed Research International, 2013, 2013, 1-10.	1.9	29
126	In Silico Prediction of Cytochrome P450-Mediated Site of Metabolism (SOM). Protein and Peptide Letters, 2013, 20, 279-289.	0.9	0

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127	Selective Synthesis of 5,6â€Dihydroindolo[1,2â€ <i>a</i>]quinoxalines and 6,7â€Dihydroindolo[2,3â€ <i>c</i>]quinolines by Orthogonal Copper and Palladium Catalysis. European Journal of Organic Chemistry, 2013, 2013, 5710-5715.	2.4	14
128	Predicting Chemical Toxicity Effects Based on Chemical-Chemical Interactions. PLoS ONE, 2013, 8, e56517.	2.5	25
129	Identifying Chemicals with Potential Therapy of HIV Based on Protein-Protein and Protein-Chemical Interaction Network. PLoS ONE, 2013, 8, e65207.	2.5	17
130	In Silico Prediction of Cytochrome P450-Mediated Site of Metabolism (SOM). Protein and Peptide Letters, 2013, 20, 279-289.	0.9	10
131	Computational Models for Predicting Interactions with Membrane Transporters. Current Medicinal Chemistry, 2013, 20, 2118-2136.	2.4	8
132	Non-Covalent Interactions with Aromatic Rings: Current Understanding and Implications for Rational Drug Design. Current Pharmaceutical Design, 2013, 19, 6522-6533.	1.9	33
133	Characterization of acetyl-CoA and propionyl-CoA carboxylases encoded by <italic>Leptospira interrogans</italic> serovar Lai: an initial biochemical study for leptospiral gluconeogenesis via anaplerotic CO ₂ assimilation. Acta Biochimica Et Biophysica Sinica. 2012. 44. 692-702.	2.0	1
134	An Investigation of the Catalytic Activity of CYP2A13*4 with Coumarin and Polymorphisms of CYP2A13 in a Chinese Han Population. Drug Metabolism and Disposition, 2012, 40, 847-851.	3.3	2
135	SOMEViz: A Web Service for Site of Metabolism Estimating and Visualizing. Protein and Peptide Letters, 2012, 19, 905-909.	0.9	2
136	Design, Synthesis, and Pharmacological Evaluation of Monocyclic Pyrimidinones as Novel Inhibitors of PDE5. Journal of Medicinal Chemistry, 2012, 55, 10540-10550.	6.4	28
137	Development of a novel class of B-RafV600E-selective inhibitors through virtual screening and hierarchical hit optimization. Organic and Biomolecular Chemistry, 2012, 10, 7402.	2.8	20
138	Estimation of Carcinogenicity Using Molecular Fragments Tree. Journal of Chemical Information and Modeling, 2012, 52, 1994-2003.	5.4	15
139	Unbinding Pathways of GW4064 from Human Farnesoid X Receptor As Revealed by Molecular Dynamics Simulations. Journal of Chemical Information and Modeling, 2012, 52, 3043-3052.	5.4	22
140	Toward understanding the molecular basis for chemical allosteric modulator design. Journal of Molecular Graphics and Modelling, 2012, 38, 324-333.	2.4	36
141	Discovery of new non-steroidal FXR ligands via a virtual screening workflow based on Phase shape and induced fit docking. Bioorganic and Medicinal Chemistry Letters, 2012, 22, 6848-6853.	2.2	22
142	Recent Advances in Neuraminidase Inhibitor Development as Antiâ€influenza Drugs. ChemMedChem, 2012, 7, 1527-1536.	3.2	62
143	Dynamic monitoring of \hat{l}^2 -cell injury with impedance and rescue by glucagon-like peptide-1. Analytical Biochemistry, 2012, 423, 61-69.	2.4	9
144	Enantioselective drug–protein interaction between mexiletine and plasma protein. Journal of Pharmacy and Pharmacology, 2012, 64, 792-801.	2.4	9

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145	(+)-Rutamarin as a Dual Inducer of Both GLUT4 Translocation and Expression Efficiently Ameliorates Glucose Homeostasis in Insulin-Resistant Mice. PLoS ONE, 2012, 7, e31811.	2.5	28
146	Knowledge-Based Scoring Functions in Drug Design: 2. Can the Knowledge Base Be Enriched?. Journal of Chemical Information and Modeling, 2011, 51, 386-397.	5.4	31
147	Knowledge-Based Scoring Functions in Drug Design: 3. A Two-Dimensional Knowledge-Based Hydrogen-Bonding Potential for the Prediction of Protein–Ligand Interactions. Journal of Chemical Information and Modeling, 2011, 51, 2994-3004.	5.4	17
148	Computational Screening for Active Compounds Targeting Protein Sequences: Methodology and Experimental Validation. Journal of Chemical Information and Modeling, 2011, 51, 2821-2828.	5.4	61
149	Catalytic Mechanism Investigation of Lysine-Specific Demethylase 1 (LSD1): A Computational Study. PLoS ONE, 2011, 6, e25444.	2.5	42
150	AST1306, A Novel Irreversible Inhibitor of the Epidermal Growth Factor Receptor 1 and 2, Exhibits Antitumor Activity Both In Vitro and In Vivo. PLoS ONE, 2011, 6, e21487.	2.5	40
151	Fragment-based prediction of skin sensitization using recursive partitioning. Journal of Computer-Aided Molecular Design, 2011, 25, 885-893.	2.9	10
152	An integrated drug-likeness study for bicyclic privileged structures: from physicochemical properties to in vitro ADME properties. Molecular Diversity, 2011, 15, 857-876.	3.9	19
153	Identification and synthesis of N′-(2-oxoindolin-3-ylidene)hydrazide derivatives against c-Met kinase. Bioorganic and Medicinal Chemistry Letters, 2011, 21, 3749-3754.	2.2	45
154	Identification of pentacyclic triterpenes derivatives as potent inhibitors against glycogen phosphorylase based on 3D-QSAR studies. European Journal of Medicinal Chemistry, 2011, 46, 2011-2021.	5.5	30
155	Molecular Basis of NDM-1, a New Antibiotic Resistance Determinant. PLoS ONE, 2011, 6, e23606.	2.5	62
156	Novel thiophene derivatives as PTP1B inhibitors with selectivity and cellular activity. Bioorganic and Medicinal Chemistry, 2010, 18, 1773-1782.	3.0	59
157	Using support vector regression coupled with the genetic algorithm for predicting acute toxicity to the fathead minnow. SAR and QSAR in Environmental Research, 2010, 21, 559-570.	2.2	32
158	Knowledge-Based Scoring Functions in Drug Design. 1. Developing a Target-Specific Method for Kinaseâ~Ligand Interactions. Journal of Chemical Information and Modeling, 2010, 50, 1378-1386.	5.4	33
159	Site of metabolism prediction for six biotransformations mediated by cytochromes P450. Bioinformatics, 2009, 25, 1251-1258.	4.1	72
160	An effective docking strategy for virtual screening based on multi-objective optimization algorithm. BMC Bioinformatics, 2009, 10, 58.	2.6	39
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162	Dynamic Mechanism of Fatty Acid Transport across Cellular Membranes through FadL: Molecular Dynamics Simulations. Journal of Physical Chemistry B, 2008, 112, 13070-13078.	2.6	10

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